# **Refine Search**

# Search Results -

| Terms   | Documents |
|---|-----------|
| plant adj transformation and lecithin adj cholesterol adj acyltransferase | 0         |

US Pre-Grant Publication Full-Text Database US Patents Full-Text Database

US OCR Full-Text Database

Database:

EPO Abstracts Database JPO Abstracts Database **Derwent World Patents Index** 

IBM Technical Disclosure Bulletins

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|     | Recall Text 👄 | Clear  |    | Interrupt    |

### Search History

DATE: Friday, December 10, 2004 Printable Copy Create Case

| Set Name<br>side by side |   | Hit Count | Set Name<br>result set |
|--------------------------|---|-----------|------------------------|
| DB=PG                    | SPB,USPT,USOC,EPAB,JPAB,DWPI; PLUR=YES; OP=OR                             |           |                        |
| <u>L10</u>               | plant adj transformation and lecithin adj cholesterol adj acyltransferase | 0         | <u>L10</u>             |
| <u>L9</u>                | plant adj sterol and lecithin adj cholesterol adj acyltransferase         | 29        | <u>L9</u>              |
| <u>L8</u>                | plant sterol and lecithin adj cholesterol adj acyltransferase             | 744189    | L8                     |
| <u>L7</u>                | L4 and plant.clm.   | 2         | <u>L7</u>              |
| <u>L6</u>                | L4 and plant  | 246       | <u>L6</u>              |
| <u>L5</u>                | L1 and (lecithin near3 acyltransferase).clm.                              | 6         | <u>L5</u>              |
| <u>L4</u>                | L1 and (lecithin near3 acyltransferase)                                   | 246       | <u>L4</u>              |
| <u>L3</u>                | L1 anf (lecithin near3 acyltransferase)                                   | 4958      | <u>L3</u>              |
| <u>L.2</u>               | L1 anf lecithin near3 acyltransferase                                     | 4958      | <u>L2</u>              |
| LI                       | acyltransferase and lecithin and cholesterol and plant                    | 343       | <u>L1</u>              |
|                          |   |           |                        |

END OF SEARCH HISTORY

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FILE 'CAPLUS' ENTERED AT 15:50:34 ON 10 DEC 2004
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COPYRIGHT (C) 2004 AMERICAN CHEMICAL SOCIETY (ACS)
=> s lecithin(w)cholesterol(w)acyltransferase
          5907 LECITHIN(W) CHOLESTEROL(W) ACYLTRANSFERASE
=> s l1 and plant
            92 L1 AND PLANT
=> duplicate remove 12
DUPLICATE PREFERENCE IS 'AGRICOLA, BIOSIS, EMBASE, CAPLUS'
KEEP DUPLICATES FROM MORE THAN ONE FILE? Y/(N):n
PROCESSING COMPLETED FOR L2
L3
             64 DUPLICATE REMOVE L2 (28 DUPLICATES REMOVED)
=> s l1 and pd <1999
'1999' NOT A VALID FIELD CODE
   2 FILES SEARCHED...
   3 FILES SEARCHED...
         4738 L1 AND PD <1999
=>
   1999
1999 IS NOT A RECOGNIZED COMMAND
The previous command name entered was not recognized by the system.
For a list of commands available to you in the current file, enter
"HELP COMMANDS" at an arrow prompt (=>).
=>
=> d 13 1-10 ti
    ANSWER 1 OF 64 CAPLUS COPYRIGHT 2004 ACS on STN
L3
ΤI
    Method for producing polyunsaturated fatty acids, lipids, and oils in
     transgenic organisms expressing fungal acyltransferases
```

- L3 ANSWER 2 OF 64 BIOSIS COPYRIGHT (c) 2004 The Thomson Corporation. on STN DUPLICATE 1
- TI Expression in yeast of a novel phospholipase A1 cDNA from Arabidopsis thaliana.
- L3 ANSWER 3 OF 64 BIOSIS COPYRIGHT (c) 2004 The Thomson Corporation. on STN DUPLICATE 2
- TI Cloning and functional characterization of a Phospholipid:Diacylglycerol acyltransferase from Arabidopsis.
- L3 ANSWER 4 OF 64 BIOSIS COPYRIGHT (c) 2004 The Thomson Corporation. on STN DUPLICATE 3
- TI Compared with acyl-CoA: Cholesterol O-acyltransferase (ACAT) 1 and

  \*\*\*lecithin\*\*\* : \*\*\*Cholesterol\*\*\* \*\*\*acyltransferase\*\*\* , ACAT2
  displays the greatest capacity to differentiate cholesterol from
  sitosterol.
- L3 ANSWER 5 OF 64 EMBASE COPYRIGHT 2004 ELSEVIER INC. ALL RIGHTS RESERVED. ON STN DUPLICATE 4
- TI Pharmacotherapy for dyslipidaemia Current therapies and future agents.
- L3 ANSWER 6 OF 64 BIOSIS COPYRIGHT (c) 2004 The Thomson Corporation. on STN DUPLICATE 5
- TI The human cholesteryl ester transfer protein I405V polymorphism is associated with plasma cholesterol concentration and its reduction by dietary phytosterol esters.
- L3 · ANSWER 7 OF 64 AGRICOLA Compiled and distributed by the National Agricultural Library of the Department of Agriculture of the United States of America. It contains copyrighted materials. All rights reserved.

  (2004) on STN
- TI Accumulation of genistein and lipophilic genistein derivatives in lipoproteins during incubation with human plasma in vitro.
- L3 ANSWER 8 OF 64 BIOSIS COPYRIGHT (c) 2004 The Thomson Corporation. on STN
- TI The seeds from Plantago ovata lower plasma lipids by altering hepatic and bile acid metabolism in guinea pigs.
- L3 ANSWER 9 OF 64 BIOSIS COPYRIGHT (c) 2004 The Thomson Corporation. on STN DUPLICATE 6
- TI Lipoprotein-associated estrogens.
- L3 ANSWER 10 OF 64 BIOSIS COPYRIGHT (c) 2004 The Thomson Corporation. on STN DUPLICATE 7
- TI Lipid lowering activity of Phyllanthus niruri in hyperlipemic rats.

### => d 13 1-4 ibib ab

L3 ANSWER 1 OF 64 CAPLUS COPYRIGHT 2004 ACS on STN

ACCESSION NUMBER:

2004:857694 CAPLUS

DOCUMENT NUMBER:

141:344590

TITLE:

Method for producing polyunsaturated fatty acids, lipids, and oils in transgenic organisms expressing

fungal acyltransferases

INVENTOR(S):

Renz, Andreas; Bauer, Joerg; Frentzen, Margit; Soezer,

Nursen; Keith, Stobart; Fraser, Thomas; Lazarus, Colin

M.; Qi, Baoxiu; Abbadi, Amine; Heinz, Ernst

PATENT ASSIGNEE(S):

University of Bristol, UK PCT Int. Appl., 270 pp.

SOURCE:

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

German

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

| PAT   | ENT        | NO. |      |     | KIN | D   | DATE     |     |     | APPL           | ICAT | ION  | . 01/ |     | Di       | ATE . |     |  |
|-------|------------|-----|------|-----|-----|-----|----------|-----|-----|----------------|------|------|-------|-----|----------|-------|-----|--|
| WO    | 2004087902 |     |      |     | A2  | -   | 20041014 |     | 1   | WO 2004-EP3224 |      |      |       |     | 20040326 |       |     |  |
|       | W:         | ΑE, | AG,  | AL, | AM, | AT, | AU,      | AZ, | BA, | BB,            | BG,  | BR,  | BW,   | BY, | ΒZ,      | CA,   | CH, |  |
|       |            | CN, | CO,  | CR, | CU, | CZ, | DE,      | DK, | DM, | DZ,            | EC,  | EE,  | EG,   | ES, | FΙ,      | GB,   | GD, |  |
|       |            | GE, | GH,  | GM, | HR, | HU, | ID,      | IL, | IN, | IS,            | JP,  | KE,  | KG,   | ΚP, | KR,      | KΖ,   | LC, |  |
|       |            | LK, | LR,  | LS, | LT, | LU, | LV,      | MA, | MD, | MG,            | MK,  | MN,  | MW,   | MX, | MZ,      | NA,   | NI, |  |
|       |            | NO, | NZ,  | OM, | PG, | PH, | PL,      | PT, | RO, | RU,            | SC,  | SD,  | SE,   | SG, | SK,      | SL,   | SY, |  |
|       |            | TJ, | TM,  | TN, | TR, | TT, | TZ,      | UA, | UG, | US,            | UZ,  | VC,  | VN,   | YU, | ZA,      | ZM,   | ZW  |  |
|       | RW:        | BW, | GH,  | GM, | KΕ, | LS, | MW,      | MZ, | SD, | SL,            | SZ,  | TZ,  | UG,   | ZM, | ZW,      | AM,   | AZ, |  |
|       |            | BY, | KG,  | KΖ, | MD, | RU, | ТJ,      | TM, | AT, | BE,            | BG,  | CH,  | CY,   | CZ, | DE,      | DK,   | EE, |  |
|       |            | ES, | FI,  | FR, | GB, | GR, | HU,      | ΙE, | IT, | LU,            | MC,  | NL,  | PL,   | PT, | RO,      | SE,   | SI, |  |
|       |            | SK, | TR,  | BF, | ВJ, | CF, | CG,      | CI, | CM, | GA,            | GN,  | GQ,  | GW,   | ML, | MR,      | NE,   | SN, |  |
|       |            | TD, | TG   |     |     |     |          |     |     |                |      |      |       |     |          |       |     |  |
| ORITY | APP        | LN. | INFO | .:  |     |     |          |     | ]   | DE 2           | 003- | 1031 | 1759  | i   | A 20     | 00303 | 331 |  |
|       |            |     |      |     |     |     |          |     |     |                |      |      |       |     | _        |       |     |  |

PRIO DE 2003-10348996 A 20031017

AΒ The invention relates to a method for the prodn. of long-chained, multiply unsatd. fatty acids in an organism, wherein nucleic acids coding for proteins with acyltransferase activity are introduced into the organism. Said nucleic acid sequences can be advantageously expressed in the organism, optionally together with other nucleic acid sequences encoding enzymes involved in the biosynthesis of fatty acids or in lipid metab. The invention also relates to a method for the prodn. of oils and/or triacylglycerides with an increased content of long-chained, multiply unsatd. fatty acids. The invention further relates to the nucleic acid sequences, vectors contq. the nucleic acid sequences, and transgenic organisms contq. the above-mentioned nucleic acid sequences or vectors. The invention addnl. relates to oils, lipids and/or fatty acids produced according to the inventive method and to the utilization thereof in feed, food, cosmetics, and pharmaceuticals. Thus, lysophosphatidic acid acyltransferase, glycerol-3-phosphate acyltransferase, diacylglyerol acyltransferase, and \*\*\*lecithin\*\*\* - \*\*\*cholesterol\*\*\*

\*\*\*acyltransferase\*\*\* of Thraustochytrium, Physcomitrella patens, Cryptothecodinium cohnii, Mortierella alpina, Shewanella hanedai, and Fusarium graminearum and the corresponding cDNAs are disclosed. Acyl CoA:lysophospholipid acyltransferase cDNAs of Caenorhabditis elegans were cloned, sequenced, and expressed in yeast, tobacco, and flax and the alteration of the lipid profile was detd. The fungal acyltransferases were expressed in A. thaliana, tobacco, flax, and rape.

L3 ANSWER 2 OF 64 BIOSIS COPYRIGHT (c) 2004 The Thomson Corporation. on STN DUPLICATE 1

ACCESSION NUMBER: DOCUMENT NUMBER:

2004:441893 BIOSIS PREV200400446784

TITLE:

Expression in yeast of a novel phospholipase Al cDNA from

Arabidopsis thaliana.

AUTHOR (S):

Noiriel, Alexandre; Benveniste, Pierre; Banas, Antoni;

Stymne, Sten; Bouvier-Nave, Pierrette [Reprint Author]

CORPORATE SOURCE: CNRSInst Biol Mol PlantesDept Isoprenoides, Inst Bot, 28

Rue Goethe, F-67083, Strasbourg, France Pierrette.Nave@bota-ulp.u-strasbg.fr

SOURCE: European Journal of Biochemistry, (September 2004) Vol.

271, No. 18, pp. 3752-3764. print.

ISSN: 0014-2956 (ISSN print).

DOCUMENT TYPE:

Article English

LANGUAGE: ENTRY DATE:

Entered STN: 17 Nov 2004

Last Updated on STN: 17 Nov 2004

AB During a search for cDNAs encoding \*\*\*plant\*\*\* sterol acyltransferases, we isolated four full-length cDNAs from Arabidopsis

thaliana that encode proteins with substantial identity with animal

\*\*\*lecithin\*\*\* : \*\*\*cholesterol\*\*\* \*\*\*acyltransferases\*\*\*

(LCATs). The expression of one of these cDNAs, AtLCAT3 (At3g03310), in
various yeast strains resulted in the doubling of the triacylglycerol
content. Furthermore, a complete lipid analysis of the transformed
wild-type yeast showed that its phospholipid content was lower than that
of the control (void plasmid-transformed) yeast whereas lysophospholipids
and free fatty acids increased. When microsomes from the

and free fatty acids increased. When microsomes from the AtLCAT3-transformed yeast were incubated with di-(1-14C)oleyl phosphatidylcholine, both the lysophospholipid and free fatty acid fractions were highly and similarly labelled, whereas the same incubation with microsomes from the control yeast produced a negligible labelling of these fractions. Moreover when microsomes from AtLCAT3-transformed yeast were incubated with either sn-1- or sn-2-(1-14C)acyl phosphatidylcholine, the distribution of the labelling between the free fatty acid and the lysophosphatidylcholine fractions strongly suggested a phospholipase A1 activity for AtLCAT3. The sn-1 specificity of this phospholipase was

confirmed by gas chromatography analysis of the hydrolysis of 1-myristoyl, 2-oleyl phosphatidylcholine. Phosphatidylethanolamine and phosphatidic acid were shown to be also hydrolysed by AtLCAT3, although less efficiently than phosphatidylcholine. Lysophospatidylcholine was a weak substrate whereas tripalmitoylglycerol and cholesteryl oleate were not hydrolysed at all. This novel A. thaliana phospholipase Al shows optimal

activity at pH 6-6.5 and 60-65 degreeC and appears to be unaffected by Ca2+. Its sequence is unrelated to all other known phospholipases. Further studies are in progress to elucidate its physiological role.

L3 ANSWER 3 OF 64 BIOSIS COPYRIGHT (c) 2004 The Thomson Corporation. on STN DUPLICATE 2

ACCESSION NUMBER: DOCUMENT NUMBER:

2004:354070 BIOSIS PREV200400354408

TITLE:

Cloning and functional characterization of a

Phospholipid:Diacylglycerol acyltransferase from

Arabidopsis.

AUTHOR(S):

Stahl, Ulf; Carlsson, Anders S. [Reprint Author]; Lenman,

Marit; Dahlqvist, Anders; Huang, Bangquan; Banas,

Walentyna; Banas, Antoni; Stymne, Sten

CORPORATE SOURCE:

Dept Crop Sci, Swedish Univ Agr Sci, S-23053, Alnarp,

Sweden

anders.carlsson@vv.slu.se

SOURCE:

Plant Physiology (Rockville), (July 2004) Vol. 135, No. 3,

pp. 1324-1335. print.

ISSN: 0032-0889 (ISSN print).

DOCUMENT TYPE:

Article

LANGUAGE:

English

ENTRY DATE:

Entered STN: 26 Aug 2004

Last Updated on STN: 26 Aug 2004

AB A new pathway for triacylglycerol biosynthesis involving a phospholipid:diacylglycerol acyltransferase (PDAT) was recently described (Dahlqvist A, Stahl U, Lenman M, Banas A, Lee M, Sandager L, Ronne H, Stymne S, (2000) Proc Natl Acad Sci USA 97: 6487-6492). The LRO1 gene that encodes the PDAT was identified in yeast (Saccharomyces cerevisiae) and shown to have homology with animal \*\*\*lecithin\*\*\*:

\*\*\*cholesterol\*\*\* \*\*\*acyltransferase\*\*\* . A search of the Arabidopsis genome database identified the protein encoded by the At5g13640 gene as the closest homolog to the yeast PDAT (28% amino acid identity). The cDNA of At5g13640 (AtPDAT gene) was overexpressed in Arabidopsis behind the cauliflower mosaic virus promoter. Microsomal preparations of roots and leaves from overexpressers had PDAT activities that correlated with expression levels of the gene, thus demonstrating that this gene encoded PDAT (AtPDAT). The AtPDAT utilized different phospholipids as acyl donor and accepted acyl groups ranging from C10 to C22. The rate of activity was highly dependent on acyl composition with highest activities for acyl groups containing several double bonds, epoxy, or hydroxy groups. The enzyme utilized both sn-positions of phosphatidylcholine but had a 3-fold preference for the sn-2 position. The fatty acid and lipid composition as well as the amounts of lipids per fresh weight in Arabidopsis \*\*\*plants\*\*\* overexpressing AtPDAT were not significantly different from the wild type. Microsomal preparations of roots from a T-DNA insertion mutant in the AtPDAT gene had barely detectable capacity to transfer acyl groups from phospholipids to added diacylglycerols. However, these microsomes were still able to carry out triacylglycerol synthesis by a diacylglycerol:diacylglycerol acyltransferase reaction at the same rate as microsomal preparations from `wild type.

L3 ANSWER 4 OF 64 BIOSIS COPYRIGHT (c) 2004 The Thomson Corporation. on STN DUPLICATE 3

ACCESSION NUMBER:

2004:72411 BIOSIS

DOCUMENT NUMBER:

PREV200400076068

TITLE:

Compared with acyl-CoA: Cholesterol O-acyltransferase (ACAT) 1 and \*\*\*lecithin\*\*\* : \*\*\*Cholesterol\*\*\*

\*\*\*acyltransferase\*\*\* , ACAT2 displays the greatest capacity to differentiate cholesterol from sitosterol.

AUTHOR(S):

Temel, Ryan E. [Reprint Author]; Gebre, Abraham K.; Parks,

John S.; Rudel, Lawrence L.

CORPORATE SOURCE:

Dept. of Pathology, Wake Forest University School of

Medicine, Hanes Bldg., 6th Floor, Winston-Salem, NC, 27157,

USA

rtemel@wfubmc.edu; lrudel@wfubmc.edu

SOURCE:

Journal of Biological Chemistry, (November 28 2003) Vol.

278, No. 48, pp. 47594-47601. print.

CODEN: JBCHA3. ISSN: 0021-9258.

DOCUMENT TYPE:

Article

LANGUAGE:

English

ENTRY DATE:

Entered STN: 4 Feb 2004

Last Updated on STN: 4 Feb 2004

The capacity of acyl-CoA:cholesterol O-acyltransferase (ACAT) 2 to differentiate cholesterol from the \*\*\*plant\*\*\* sterol, sitosterol, was compared with that of the sterol esterifying enzymes, ACAT1 and \*\*\*lecithin\*\*\* : \*\*\*cholesterol\*\*\* \*\*\*acyltransferase\*\*\* (LCAT).

Cholesterol-loaded microsomes from transfected cells containing either ACAT1 or ACAT2 exhibited significantly more ACAT activity than their sitosterol-loaded counterparts. In sitosterolloaded microsomes, both ACAT1 and ACAT2 were able to esterify sitosterol albeit with lower efficiencies than cholesterol. The mass ratios of cholesterol ester to sitosterol ester formed by ACAT1 and ACAT2 were 1.6 and 7.2, respectively. Compared with ACAT1, ACAT2 selectively esterified cholesterol even when sitosterol was loaded into the microsomes. To further characterize the difference in sterol specificity, ACAT1 and ACAT2 were compared in intact cells loaded with either cholesterol or sitosterol. Despite a lower level of ACAT activity, the ACAT1-expressing cells esterified 4-fold more sitosterol than the ACAT2 cells. The data showed that compared with ACAT1, ACAT2 displayed significantly greater selectively for cholesterol compared with sitosterol. The plasma cholesterol esterification enzyme \*\*\*lecithin\*\*\* : \*\*\*cholesterol\*\*\* \*\*\*acyltransferase\*\*\* compared. With recombinant high density lipoprotein particles, the esterification rate of cholesterol by LCAT was only 15% greater than for sitosterol. Thus, LCAT was able to efficiently esterify both cholesterol and sitosterol. In contrast, ACAT2 demonstrated a strong preference for cholesterol rather than sitosterol. This sterol selectivity by ACAT2 may reflect a role in the sorting of dietary sterols during their absorption by the intestine in vivo.

#### => d 13 5-10 ibib

ANSWER 5 OF 64 EMBASE COPYRIGHT 2004 ELSEVIER INC. ALL RIGHTS RESERVED. on STN DUPLICATE 4

ACCESSION NUMBER:

2003463371 EMBASE

TITLE:

Pharmacotherapy for dyslipidaemia - Current therapies and

future agents.

AUTHOR:

Bays H.; Stein E.A.

CORPORATE SOURCE:

Dr. H. Bays, L-MARC Research Center, 3288 Illinois Avenue,

Louisville, KY 40213, United States. HBaysMD@aol.com

SOURCE:

Expert Opinion on Pharmacotherapy, (2003) 4/11 (1901-1938).

Refs: 225

ISSN: 1465-6566 CODEN: EOPHF7

COUNTRY:

United Kingdom

DOCUMENT TYPE:

Journal; General Review

FILE SEGMENT:

018 Cardiovascular Diseases and Cardiovascular Surgery

037 Drug Literature Index 038 Adverse Reactions Titles

LANGUAGE:

English

SUMMARY LANGUAGE: English

ANSWER 6 OF 64 BIOSIS COPYRIGHT (c) 2004 The Thomson Corporation. L3DUPLICATE 5

ACCESSION NUMBER:

2003:393370 BIOSIS

DOCUMENT NUMBER:

PREV200300393370

TITLE:

The human cholesteryl ester transfer protein I405V polymorphism is associated with plasma cholesterol concentration and its reduction by dietary phytosterol

AUTHOR(S):

Lottenberg, Ana M. [Reprint Author]; Nunes, Valeria S.; Nakandakare, Edna R.; Neves, Monica; Bernik, Marcia; Lagrost, Laurent; dos Santos, Jose E.; Quintao, Eder

CORPORATE SOURCE:

Lipid Laboratory, Medical School, University of Sao Paulo,

LIM10, Sao Paulo, Brazil

lipideq@usp.br

SOURCE:

Journal of Nutrition, (June 2003) Vol. 133, No. 6, pp.

1800-1805. print.

ISSN: 0022-3166 (ISSN print).

DOCUMENT TYPE:

Article

LANGUAGE:

English

ENTRY DATE:

Entered STN: 27 Aug 2003

Last Updated on STN: 27 Aug 2003

L3ANSWER 7 OF 64 AGRICOLA Compiled and distributed by the National Agricultural Library of the Department of Agriculture of the United States of America. It contains copyrighted materials. All rights reserved. (2004) on STN

ACCESSION NUMBER:

2003:30671 AGRICOLA

DOCUMENT NUMBER:

IND23326146

TITLE:

Accumulation of genistein and lipophilic genistein

derivatives in lipoproteins during incubation with

human plasma in vitro.

AUTHOR (S):

Kaamanen, M.; Adlercreutz, H.; Jauhiainen, M.;

Tikkanen, M.J.

AVAILABILITY:

DNAL (381 B522)

SOURCE:

Biochimica et biophysica acta = International journal of biochemistry and biophysics, Mar 17, 2003. Vol.

1631, No. 2. p. 147-152

Publisher: Amsterdam : Elsevier Science B.V.

CODEN: BBACAQ; ISSN: 0006-3002

NOTE:

Includes references

PUB. COUNTRY:

DOCUMENT TYPE:

Article

FILE SEGMENT:

Non-U.S. Imprint other than FAO

LANGUAGE:

English

ANSWER 8 OF 64 BIOSIS COPYRIGHT (c) 2004 The Thomson Corporation. on 1.3

STN

ACCESSION NUMBER:

2002:362034 BIOSIS

Netherlands

DOCUMENT NUMBER:

AUTHOR (S):

PREV200200362034

TITLE:

The seeds from Plantago ovata lower plasma lipids by

altering hepatic and bile acid metabolism in guinea pigs. Romero, Ana Lourdes [Reprint author]; West, Kristy L.;

Zern, Tosca; Fernandez, Maria Luz

CORPORATE SOURCE:

Department of Food Science, University of Sonora,

Hermosillo, SON, Mexico

maria-luz.fernandez@uconn.edu

SOURCE:

Journal of Nutrition, (June, 2002) Vol. 132, No. 6, pp.

1194-1198. print.

CODEN: JONUAI. ISSN: 0022-3166.

DOCUMENT TYPE:

Article English

LANGUAGE: ENTRY DATE:

Entered STN: 26 Jun 2002

Last Updated on STN: 26 Jun 2002

ANSWER 9 OF 64 BIOSIS COPYRIGHT (c) 2004 The Thomson Corporation. L3STN

DUPLICATE 6

ACCESSION NUMBER:

2003:24021 BIOSIS PREV200300024021

DOCUMENT NUMBER: TITLE:

Lipoprotein-associated estrogens.

AUTHOR(S): Tikkanen, Matti J. [Reprint Author]; Vihma, Veera;

Jauhiainen, Matti; Hockerstedt, Anna; Helisten,

Hannamaarit; Kaamanen, Maija

CORPORATE SOURCE: Department of Medicine, Division of Cardiology, Helsinki

University Central Hospital, 00290, Helsinki, Finland

matti.tikkanen@hus.fi

SOURCE: Cardiovascular Research, (November 2002) Vol. 56, No. 2,

pp. 184-188. print.

CODEN: CVREAU. ISSN: 0008-6363.

DOCUMENT TYPE: Article

General Review; (Literature Review)

LANGUAGE:

English

ENTRY DATE: Entered STN: 1 Jan 2003

Last Updated on STN: 1 Jan 2003

ANSWER 10 OF 64 BIOSIS COPYRIGHT (c) 2004 The Thomson Corporation.

DUPLICATE 7

ACCESSION NUMBER:

2002:541156 BIOSIS

DOCUMENT NUMBER: TITLE:

PREV200200541156 Lipid lowering activity of Phyllanthus niruri in

hyperlipemic rats.

AUTHOR(S): Khanna, A. K.; Rizvi, F.; Chander, R. [Reprint author]

Division of Biochemistry, Central Drug Research Institute, CORPORATE SOURCE:

Lucknow, 226001, India

SOURCE:

Journal of Ethnopharmacology, (September, 2002) Vol. 82,

No. 1, pp. 19-22. print.

CODEN: JOETD7. ISSN: 0378-8741.

DOCUMENT TYPE:

Article English

LANGUAGE: ENTRY DATE:

Entered STN: 16 Oct 2002

Last Updated on STN: 16 Oct 2002

=> d 13 11-20 ibib

ANSWER 11 OF 64 CAPLUS COPYRIGHT 2004 ACS on STN

ACCESSION NUMBER:

2001:168132 CAPLUS

DOCUMENT NUMBER:

134:218021

TITLE:

Nucleic acids encoding \*\*\*plant\*\*\*

acyltransferases and their use to modify sterol

composition

INVENTOR(S): Lassner, Michael; Van Eenennaam, Alison

PATENT ASSIGNEE(S):

Monsanto Company, USA

SOURCE:

PCT Int. Appl., 127 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

| PATENT NO.       |                  |    |     | KIN | D   | DATE                     |      |            | APPLICATION NO. |     |     |     |     |     | DATE     |     |     |     |
|------------------|------------------|----|-----|-----|-----|--------------------------|------|------------|-----------------|-----|-----|-----|-----|-----|----------|-----|-----|-----|
|                  |                  |    |     |     |     |                          | _    | <b>-</b> - |                 |     |     |     |     |     |          |     |     |     |
| WO 2001016308 A2 |                  |    |     |     |     | 20010308 WO 2000-US23863 |      |            |                 |     |     |     |     | 2   | 20000830 |     |     |     |
|                  | WO 2001016308 A3 |    |     |     |     |                          | 2002 | 0117       |                 |     |     |     |     |     |          |     |     |     |
|                  |                  | W: | ΑE, | AG, | AL, | AM,                      | AT,  | AU,        | AZ,             | BA, | BB, | BG, | BR, | BY, | BZ,      | CA, | CH, | CN, |
|                  |                  |    |     |     |     |                          |      | DM,        |                 |     |     |     |     |     |          |     |     |     |
|                  |                  |    | HU, | ID, | ΙL, | IN,                      | IS,  | JP,        | KE,             | KG, | ΚP, | KR, | KZ, | LC, | LK,      | LR, | LS, | LT, |

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LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU,
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             YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
         RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY,
             DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ,
             CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
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                                 20020507
                                             BR 2000-14154
                           Α
                                                                     20000830
     EP 1210417
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                                             EP 2000-959644
                                                                     20000830
         R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
             IE, SI, LT, LV, FI, RO, MK, CY, AL
     JP 2003508052
                          T2
                                 20030304
                                             JP 2001-520855
                                                                     20000830
     ZA 2002001410
                           Α
                                 20030606
                                             ZA 2002-1410
                                                                     20020219
PRIORITY APPLN. INFO.:
                                             US 1999-152493P
                                                                     19990830
                                             WO 2000-US23863
                                                                    20000830
     ANSWER 12 OF 64 BIOSIS COPYRIGHT (c) 2004 The Thomson Corporation.
     STN
                                                         DUPLICATE 8
ACCESSION NUMBER:
                    2001:414984 BIOSIS
DOCUMENT NUMBER:
                    PREV200100414984
TITLE:
                    Effect of administration with the extract of Gymnema
                    sylvestre R. Br leaves on lipid metabolism in rats.
                    Shigematsu, Norihiro [Reprint author]; Asano, Ryuji;
AUTHOR(S):
                    Shimosaka, Makoto; Okazaki, Mitsuo
                    Biosci. Textile Technol., Shinshu University, 3-15-1
CORPORATE SOURCE:
                    Tokida, Ueda, Nagano, 386-8567, Japan
                    noshigematsu@fancl.co.jp
                    Biological and Pharmaceutical Bulletin, (June, 2001) Vol.
SOURCE:
                    24, No. 6, pp. 713-717. print.
                    ISSN: 0918-6158.
DOCUMENT TYPE:
                    Article
LANGUAGE:
                    English
ENTRY DATE:
                    Entered STN: 29 Aug 2001
                    Last Updated on STN: 22 Feb 2002
     ANSWER 13 OF 64 BIOSIS COPYRIGHT (c) 2004 The Thomson Corporation. on
     STN
ACCESSION NUMBER:
                    2001:246284 BIOSIS
DOCUMENT NUMBER:
                    PREV200100246284
TITLE:
                    The effect of estrogen and
                                                  ***plant***
                                                                estrogens on
                    lipoproteins.
                    Tikkanen, M. J. [Reprint author]; Adlercreutz, H. [Reprint
AUTHOR (S):
                    author]; Helisten, H. [Reprint author]; Hockerstedt, A.
                    [Reprint author]; Jauhiainen, M. [Reprint author];
                    Kaamanen, M. [Reprint author]; Tiitinen, A. [Reprint
                    author]; Wahala, K. [Reprint author]
CORPORATE SOURCE:
                    Dept. of Medicine, University of Helsinki, Helsinki,
                    Finland
SOURCE:
                    Pfluegers Archiv European Journal of Physiology, (2001)
                    Vol. 441, No. 6 Supplement, pp. R121. print.
                    Meeting Info.: Joint Congress of the Scandinavian and the
                    German Physiological Societies. Berlin, Germany. March
                    10-13, 2001.
                    CODEN: PFLABK. ISSN: 0031-6768.
DOCUMENT TYPE:
                    Conference; (Meeting)
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Conference; Abstract; (Meeting Abstract)

English

LANGUAGE:

ENTRY DATE:

Entered STN: 23 May 2001

Last Updated on STN: 19 Feb 2002

L3 ANSWER 14 OF 64 CAPLUS COPYRIGHT 2004 ACS on STN

ACCESSION NUMBER:

2000:384442 CAPLUS

DOCUMENT NUMBER:

133:27387

TITLE:

Polynucleotides (cDNA) and polypeptides of

\*\*\*plant\*\*\* \*\*\*lecithin\*\*\* \*\*\*cholesterol\*\*\*

\*\*\*acyltransferase\*\*\* sequence homologs, sequences

and biological uses thereof

INVENTOR(S):

Cahoon, Rebecca E.; Kinney, Anthony J.; Sakai, Hajime;

Shen, Jennie Bih-jien; Butler, Karlene H.; Saylor,

James J.

PATENT ASSIGNEE(S):

E. I. Du Pont de Nemours & Co., USA

PCT Int. Appl., 49 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

SOURCE:

English

1

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

| PA       | PATENT NO. |            |      |     |     | KIND DATE |             |      | APPLICATION NO. |          |              |       |     |             | DATE |       |     |  |
|----------|------------|------------|------|-----|-----|-----------|-------------|------|-----------------|----------|--------------|-------|-----|-------------|------|-------|-----|--|
|          |            | <b>-</b> - |      |     |     | _         | <del></del> |      |                 | <b>-</b> | - <b>-</b> - |       |     | - <u></u> - | _    |       |     |  |
| WO       | 2000       | 0327       | 91   |     | A2  |           | 2000        | 0608 |                 | WO 1     | 999-         | US28. | 586 |             | 1:   | 9991: | 202 |  |
| WO       | 2000       | 0327       | 91   |     | А3  |           | 2000        | 0914 |                 |          |              |       |     |             |      |       |     |  |
|          | W:         | ΑE,        | AL,  | AU, | BA, | BB,       | BG,         | BR,  | CA,             | CN,      | CR,          | CU,   | CZ, | DM,         | EE,  | GD,   | GE, |  |
|          |            |            |      |     |     |           | IS,         |      |                 |          |              |       |     |             |      |       |     |  |
|          |            |            |      |     |     |           | RO,         |      |                 |          |              |       |     |             |      |       |     |  |
|          |            | YU,        | ZA,  | AM, | ΑZ, | BY,       | KG,         | KZ,  | MD,             | RU,      | TJ,          | TM    |     |             |      |       |     |  |
|          | RW:        | GH,        | GM,  | KE, | LS, | MW,       | SD,         | SL,  | SZ,             | TZ,      | UG,          | ZW,   | AT, | BE,         | CH,  | CY,   | DE, |  |
|          |            |            |      |     |     |           | GR,         |      |                 |          |              |       |     |             |      |       |     |  |
|          |            | CG,        | CI,  | CM, | GA, | GN,       | GW,         | ML,  | MR,             | NE,      | SN,          | TD,   | TG  |             |      |       |     |  |
| PRIORITY | Y APP      | LN.        | INFO | . : |     |           |             |      | 1               | US 1     | 998-         | 1107  | 82P | . ]         | P 19 | 9981: | 203 |  |

L3 ANSWER 15 OF 64 BIOSIS COPYRIGHT (c) 2004 The Thomson Corporation. on STN DUPLICATE 9

ACCESSION NUMBER:

2000:345817 BIOSIS

DOCUMENT NUMBER:

PREV200000345817

TITLE:

Phospholipid:diacylglycerol acyltransferase: An enzyme that

catalyzes the acyl-CoA-independent formation of

triacylglycerol in yeast and \*\*\*plants\*\*\*

AUTHOR(S):

Dahlqvist, Anders [Reprint author]; Stahl, Ulf; Lenman, Marit; Banas, Antoni; Lee, Michael; Sandager, Line; Ronne,

Hans; Stymne, Sten

CORPORATE SOURCE:

Scandinavian Biotechnology Research (ScanBi) AB, Herman

Ehles Vag 2, S-26831, Svalov, Sweden

SOURCE:

Proceedings of the National Academy of Sciences of the United States of America, (June 6, 2000) Vol. 97, No. 12, pp. 6487-6492. print.

CODEN: PNASA6. ISSN: 0027-8424.

DOCUMENT TYPE:

Article

LANGUAGE:

English

ENTRY DATE:

Entered STN: 16 Aug 2000

Last Updated on STN: 7 Jan 2002

L3 ANSWER 16 OF 64 BIOSIS COPYRIGHT (c) 2004 The Thomson Corporation. on STN DUPLICATE 10

ACCESSION NUMBER:

2000:341797 BIOSIS

DOCUMENT NUMBER:

PREV200000341797

TITLE:

Pharmacological properties of \*\*\*plant\*\*\* sterols: In

vivo and in vitro observations.

AUTHOR (S):

Moghadasian, Mohammed H. [Reprint author]

CORPORATE SOURCE:

Healthy Heart Program and Department of Pathology and

Laboratory Medicine, St. Paul's Hospital and the University of British Columbia, 180-1081 Burrard St., Vancouver, BC,

V6Z 1Y6, Canada

SOURCE:

Life Sciences, (June 30, 2000) Vol. 67, No. 6, pp. 605-615.

print.

CODEN: LIFSAK. ISSN: 0024-3205.

DOCUMENT TYPE:

Article

General Review; (Literature Review)

LANGUAGE:

English

ENTRY DATE:

Entered STN: 10 Aug 2000

Last Updated on STN: 7 Jan 2002

L3 ANSWER 17 OF 64 BIOSIS COPYRIGHT (c) 2004 The Thomson Corporation. on STN

SIN SIN

ACCESSION NUMBER:

2000:302153 BIOSIS

DOCUMENT NUMBER:

PREV200000302153

TITLE:

Astragalus mongholicus and Angelica sinensis compound

alleviates nephrotic hyperlipidemia in rats.

AUTHOR(S):

Li Jingzi [Reprint author]; Yu Lei [Reprint author]; Li Ningjun [Reprint author]; Wang Haiyan [Reprint author]

CORPORATE SOURCE:

Department of Nephrology, Research Institute of Nephrology,

First Hospital, Beijing Medical University, Beijing,

100034, China

SOURCE:

Chinese Medical Journal (English Edition), (April, 2000)

Vol. 113, No. 4, pp. 310-314. print.

CODEN: CMJODS. ISSN: 0366-6999.

DOCUMENT TYPE:

LANGUAGE:

Article English

ENTRY DATE:

Entered STN: 12 Jul 2000

Last Updated on STN: 7 Jan 2002

L3 ANSWER 18 OF 64 BIOSIS COPYRIGHT (c) 2004 The Thomson Corporation. on STN

ACCESSION NUMBER:

1999:475710 BIOSIS

DOCUMENT NUMBER:

PREV199900475710

TITLE:

Effect of inclusion of cashew globulin (Anacardium

occidentale) to a casein diet on lipid parameters in rats.

AUTHOR(S): CORPORATE SOURCE: Prabha, S. P. S.; Rajamohan, T. [Reprint author] Department of Biochemistry, University of Kerala,

Kariavattom, Trivandrum, KER, 695 581, India

SOURCE:

Plant Foods for Human Nutrition (Dordrecht), (1998) Vol.

53, No. 1, pp. 83-92. print. CODEN: PFHNE8. ISSN: 0921-9668.

DOCUMENT TYPE:

Article

LANGUAGE:

English

ENTRY DATE:

Entered STN: 9 Nov 1999

Last Updated on STN: 9 Nov 1999

L3 ANSWER 19 OF 64 BIOSIS COPYRIGHT (c) 2004 The Thomson Corporation. on STN

ACCESSION NUMBER: 1998:308035 BIOSIS

DOCUMENT NUMBER:

PREV199800308035

TITLE:

Hypolipidemic effect of flavonoids from Solanum melongena.

AUTHOR(S):

Sudheesh, S.; Presannakumar, G.; Vijayakumar, S.;

Vijayalakshmi, N. R. [Reprint author]

CORPORATE SOURCE:

Dep. Biochemistry, Univ. Kerala, Kariavattom,

Thiruvananthapuram-695 581, India

SOURCE:

Plant Foods for Human Nutrition (Dordrecht), (1997) Vol.

51, No. 4, pp. 321-330. print. CODEN: PFHNE8. ISSN: 0921-9668.

DOCUMENT TYPE:

Article

LANGUAGE:

English

ENTRY DATE:

Entered STN: 15 Jul 1998

Last Updated on STN: 13 Aug 1998

L3 ANSWER 20 OF 64 BIOSIS COPYRIGHT (c) 2004 The Thomson Corporation. on

STN

DUPLICATE 11

ACCESSION NUMBER:

1997:387450 BIOSIS

DOCUMENT NUMBER:

1997:307430 BIOSI

TITLE:

PREV199799686653
The cholesterol-raising diterpenes from coffee beans

increase serum lipid transfer protein activity levels in

humans.

AUTHOR(S):

Van Tol, Arie [Reprint author]; Urgert, Rob; De

Jong-Caesar, Ruth; Van Gent, Teus; Scheek, Leo M.; De Roos,

Baukje; Katan, Martijn B.

CORPORATE SOURCE:

Dep. Biochem., Cardiovascular Res. Inst., Fac. Med. Health

Sciences, Erasmus Univ. Rotterdam, P.O. Box 1738, 3000 DR

Rotterdam, Netherlands

SOURCE:

Atherosclerosis, (1997) Vol. 132, No. 2, pp. 251-254.

CODEN: ATHSBL. ISSN: 0021-9150.

DOCUMENT TYPE:

Article

LANGUAGE:

English

ENTRY DATE:

Entered STN: 10 Sep 1997

Last Updated on STN: 10 Sep 1997

## => d 13 18-20 ab

L3 ANSWER 18 OF 64 BIOSIS COPYRIGHT (c) 2004 The Thomson Corporation. on STN

The effect of inclusion of cashew globulin to a casein diet on lipid AR metabolism was studied in rats fed diets with two levels of cashew globulin meal. Inclusion of cashew globulin to a casein diet produced lower levels of total cholesterol, triacylglycerol and phospholipids in the serum and tissues and lower levels of serum lipoprotein cholesterol. There was decreased cholesterogenesis in the liver as evidenced by decreased activity of HMG CoA reductase and decreased release of lipoproteins into circulation. Rats fed cashew globulin along with casein also showed higher activity of LPL in the heart and adipose tissue and higher activity of LCAT. Increased hepatic diversion of cholesterol to bile acid synthesis and increased excretion of bile acids and sterols were also observed in these groups. Activity of glucose-6-phosphate dehydrogenase and malic enzyme was decreased in rats fed cashew globulin along with casein. This study demonstrates that cashew globulins included in the diet of rats are able to alter lipid metabolism which results in lower levels of lipid parameters in the serum and tissues.

STN

AB Flavonoids extracted from the fruits of Solanum melongena (Brinjal) orally administered at a dose of 1 mg/100 g BW/day showed significant hypolipidemic action in normal and cholesterol fed rats. HMG CoA reductase activity was found to be enhanced, while activities of glucose-6-phosphate dehydrogenase and malate dehydrogenase were significantly reduced. Activities of lipoprotein lipase and plasma LCAT showed significant enhancement. A significant increase in the concentrations of hepatic and fecal bile acids and fecal neutral sterols was also observed indicating a higher rate of degradation of cholesterol.

L3 ANSWER 20 OF 64 BIOSIS COPYRIGHT (c) 2004 The Thomson Corporation. on STN DUPLICATE 11

AB Cafestol and kahweol-diterpenes present in unfiltered coffee - strongly raise serum VLDL and LDL cholesterol and slightly reduce HDL cholesterol in humans. The mechanism of action is unknown. We determined whether the coffee diterpenes may affect lipoprotein metabolism via effects on lipid transfer proteins and \*\*\*lecithin\*\*\* : \*\*\*cholesterol\*\*\*

\*\*\*acyltransferase\*\*\* in a randomized, double-blind cross-over study with 10 healthy male volunteers. Either cafestol (61-64 mg,/day) or a mixture of cafestol (60 mg/day) and kahweol (48-54 mg/day) was given for 28 days. Serum activity levels of cholesterylester transfer protein, phospholipid transfer protein and \*\*\*lecithin\*\*\* : \*\*\*cholesterol\*\*\*

\*\*\*acyltransferase\*\*\* were measured using exogenous substrate assays. Relative to baseline values, cafestol raised the mean ( +- S.D.) activity of cholesterylester transfer protein by 18 +- 12% and of phospholipid transfer protein by 21 +- 14% (both P lt 0.001). Relative to cafestol alone, kahweol had no significant additional effects. \*\*\*Lecithin\*\*\*

### => d 13 21-25 ibib ab

L3 ANSWER 21 OF 64 BIOSIS COPYRIGHT (c) 2004 The Thomson Corporation. on STN

ACCESSION NUMBER:

1997:385367 BIOSIS

DOCUMENT NUMBER:

PREV199799684570

TITLE:

Immobilized Cratylia mollis lectin as a potential matrix to isolate plasma glycoproteins, including \*\*\*lecithin\*\*\* -

AUTHOR(S):

Lima, Vera L. M.; Correia, Maria T. S.; Cechinel, Yeda M. N.; Sampaio, Claudio A. M.; Owen, James S.; Coelho, Luana

C. B. B.

CORPORATE SOURCE:

Univ. Dep. Med., Royal Free Hosp. Sch. Med., Rowland Hill

St., London NW3 2PF, UK

SOURCE:

Carbohydrate Polymers, (1997) Vol. 33, No. 1, pp. 27-32.

CODEN: CAPOD8. ISSN: 0144-8617.

DOCUMENT TYPE:

Article English

LANGUAGE: ENTRY DATE:

Entered STN: 10 Sep 1997

Last Updated on STN: 10 Sep 1997

AB A crude seed extract from the native Brazilian forage, Cratylia mollis Mart., and its purified lectin (termed Cra), were found to precipitate glycoproteins from serum. An affinity column of Cra lectin coupled to

Sepharose CL-4B was prepared and its ability to isolate qlycoproteins from human plasma compared to that of a commercial immobilized lectin, Concanavalin (Con) A-Sepharose. Although both lectins are of the alpha-D-mannose/alpha-D-glucose binding class, clear differences in the type and amount of serum glycoproteins adsorbed were seen on analysis by denaturing polyacrylamide gel electrophoresis. Similarly, when a semipurified preparation of the plasma glycoprotein, \*\*\*lecithin\*\*\* \*\*\*cholesterol\*\*\* \*\*\*acyltransferase\*\*\* (LCAT, EC 2.3.1.43) was applied to the columns some differences were evident; most LCAT was not retained by either matrix but when the bound fractions were eluted and analyzed electrophoretically the LCAT isolated by the Cra-Sepharose column was much purer. These findings suggest that immobilized Cra lectin has the potential for use in studies both to isolate and to characterize

ANSWER 22 OF 64 EMBASE COPYRIGHT 2004 ELSEVIER INC. ALL RIGHTS RESERVED.

on STN

ACCESSION NUMBER: 97043150 EMBASE

certain serum glycoproteins.

DOCUMENT NUMBER: 1997043150

TITLE: Terminalia arjuna: An ayurvedic cardiotonic, regulates

lipid metabolism in hyperlipaemic rats.

Khanna A.K.; Chander R.; Kapoor N.K. AUTHOR:

R. Chander, Instituto de Farmacologia, Facultad de Ciencias CORPORATE SOURCE:

Veterinarias, Universidad Austral de Chile, P.O. Box 567,

Valdivia, Chile

SOURCE: Phytotherapy Research, (1996) 10/8 (663-665).

Refs: 19

ISSN: 0951-418X CODEN: PHYREH

COUNTRY: United Kingdom DOCUMENT TYPE: Journal; Article

FILE SEGMENT: 018 Cardiovascular Diseases and Cardiovascular Surgery

029 Clinical Biochemistry

030 Pharmacology

037 Drug Literature Index

LANGUAGE: English SUMMARY LANGUAGE: English

The lipid lowering action of the bark powder of Terminalia arjuna (T. arjuna) has been studied in triton and cholesterol fed rats. Serum lipids were found to be lowered by T. arjuna (100 mg/kg, b.w.) in triton induced hyperlipaemia. Chronic feeding of this powder (100 mg/kg, b.w., p.o.) in animals simultaneously fed with cholesterol (25 mg/kg, b.w.) for 30 days, caused lowering in lipids and protein levels of .beta.-lipoproteins followed by an increase in high density lipoprotein-cholesterol compared with the cholesterol fed groups. T. arjuna alters lipolytic activities in plasma, liver, heart and adipose tissues of hyperlipaemic rats. The lipid lowering action of this natural product is mediated through inhibition of hepatic cholesterol biosynthesis, increased faecal bile acid excretion and enhanced plasma \*\*\*lecithin\*\*\* : \*\*\*cholesterol\*\*\*

\*\*\*acyltransferase\*\*\* activity and stimulation of receptor mediated catabolism of low density lipoprotein.

L3ANSWER 23 OF 64 EMBASE COPYRIGHT 2004 ELSEVIER INC. ALL RIGHTS RESERVED. on STN

ACCESSION NUMBER: 96314582 EMBASE

DOCUMENT NUMBER: 1996314582

TITLE: Lipid lowering activity of guggulsterone from Commiphora

mukul in hyperlipaemic rats.

AUTHOR:

Chander R.; Khanna A.K.; Kapoor N.K.

CORPORATE SOURCE:

Division of Biochemistry, Central Drug Research Institute,

PO Box 173, Lucknow 226 001, India

SOURCE:

Phytotherapy Research, (1996) 10/6 (508-511).

ISSN: 0951-418X CODEN: PHYREH

COUNTRY:

United Kingdom Journal; Article

DOCUMENT TYPE: FILE SEGMENT:

018 Cardiovascular Diseases and Cardiovascular Surgery

Clinical Biochemistry 029 048 Gastroenterology

03.0 Pharmacology

037 Drug Literature Index

LANGUAGE:

English

SUMMARY LANGUAGE:

English

The lipid lowering action of guggulsterone, the active constituent of guggulipid, has been studied in triton and cholesterol fed hyperlipaemic rats. Serum lipids were found to be lowered by guggulsterone (50 mg/kg, b.w.) in triton WR-1339 induced hyperlipaemia. Chronic feeding of this drug (5 mg/kg, b.w.) in animals simultaneously fed with cholesterol (25 mg/kg, b.w.) for 30 days, caused lowering in the lipid and apoprotein levels of very low density and low density lipoproteins in experimental animals. Guggulsterone activates lipolytic enzymes in plasma and liver as well as stimulated receptor mediated catabolism of low density lipoprotein. The hypolipidaemic activity of this drug is mediated through inhibition of hepatic cholesterol biosynthesis, increased faecal bile acid excretion and enhanced plasma \*\*\*lecithin\*\*\* : \*\*\*cholesterol\*\*\* \*\*\*acyltransferase\*\*\* activity.

ANSWER 24 OF 64 BIOSIS COPYRIGHT (c) 2004 The Thomson Corporation. L3 STN DUPLICATE 12

ACCESSION NUMBER:

1996:426281 BIOSIS PREV199699157337

DOCUMENT NUMBER: TITLE:

Treatment of severe hypercholesterolemia with a combination

of beta-sitosterol and lovastatin.

AUTHOR (S):

Richter, Werner O. [Reprint author]; Geiss, Hans C.:

Soennichsen, Andreas C.; Schwandt, Peter

CORPORATE SOURCE:

Klin. Grosshadern, Dep. Internal Med. II, Marchioninistr.

15, D-81377 Munich, Germany

SOURCE:

Current Therapeutic Research, (1996) Vol. 57, No. 7, pp.

497-505.

CODEN: CTCEA9. ISSN: 0011-393X.

DOCUMENT TYPE:

Article

LANGUAGE:

English

ENTRY DATE:

Entered STN: 26 Sep 1996

Last Updated on STN: 26 Sep 1996

The objective of this study was to determine whether adding the AB sterol beta-sitosterol to a lipid-lowering treatment

regimen

of lovastatin further decreases low-density lipoprotein (LDL) cholesterol. Thirty patients (16 men, 14 women) with a mean age of 45 +- 13 years, LDL cholesterol levels between 5.89 and 12.26 mmol/L, and triglyceride levels lt 2.82 mmol/L were randomly assigned to one of two study groups: group  ${\tt L}$ (n = 15), which received lovastatin alone, and group LS (n = 15), which received lovastatin and beta-sitosterol. All patients were first treated for 16 weeks with the maximally tolerable dose of lovastatin. Beta-sitosterol 6 g/d was then added to the treatment regimen of group LS for 12 weeks, while group L continued with lovastatin alone. In the

beta-sitosterol group, mean LDL cholesterol decreased by an additional 12.8% to 15.1%, a significant difference from the corresponding change in the group receiving lovastatin alone. After discontinuation of beta-sitosterol, LDL cholesterol increased again. The decrease in LDL and total cholesterol before the addition of beta-sitosterol was comparable in the two groups: the mean reduction in LDL cholesterol was 30.4% in group L and 25.8% in group LS; total cholesterol was decreased by 24.2% in group L and 21.1% in group LS. After the addition of beta-sitosterol, the total decrease in LDL cholesterol was 35.3% to 37.1%; the decrease in total cholesterol was 27.3% to 29.2%. No significant changes were observed in other lipid variables such as high-density lipoprotein cholesterol, lipoprotein(a), very-low-density lipoprotein triglycerides and cholesterol, apolipoprotein A-I, and \*\*\*lecithin\*\*\* - \*\*\*cholesterol\*\*\* \*\*\*acyltransferase\*\*\* activity.

L3 ANSWER 25 OF 64 BIOSIS COPYRIGHT (c) 2004 The Thomson Corporation. on

ACCESSION NUMBER:

1996:370437 BIOSIS

DOCUMENT NUMBER:

PREV199699092793

TITLE:

AUTHOR (S):

Epi-cochlioquinone A, a novel acyl-CoA:cholesterol

acyltransferase inhibitor produced by Stachybotrys bisbyi. Fujioka, Tomoyuki [Reprint author]; Yao, Keiko; Hamano,

Kiyoshi; Hosoya, Tsuyoshi; Kagaski, Takeshi; Furukawa, Yoji; Haruyama, Hideyuki; Sato, Sadao; Koga, Teiichiro;

Tsujita, Yoshio

CORPORATE SOURCE:

Pharmacol. Molecular Biol. Res. Lab., Sankyo Co. Ltd.,

1-2-58 Hiromachi, Shinagawa-ku, Tokyo 140, Japan

SOURCE: Journal of Antibiotics (Tokyo), (1996) Vol. 49, No. 5, pp. 409-413.

CODEN: JANTAJ. ISSN: 0021-8820.

DOCUMENT TYPE:

Article

LANGUAGE:

English

ENTRY DATE:

Entered STN: 14 Aug 1996

Last Updated on STN: 15 Aug 1996

AB A novel acyl-CoA: cholesterol acyltransferase (ACAT) inhibitor, designated epi-cochlioquinone A has been isolated from the fermentation broth of Stachybotrys bisbyi SANK 17777. The molecular formula, physicochemical properties, NMR spectroscopic analysis and X-ray crystallographic analysis revealed that this compound was a stereoisomer of cochlioquinone A, which has been previously reported as a nematocidal agent. It inhibited ACAT activity in an enzyme assay using rat liver microsomes with an IC-50 value of 1.7 mu-M. However, it showed about 10-fold less potent inhibitory effect on plasma \*\*\*lecithin\*\*\* \*\*\*cholesterol\*\*\*

\*\*\*acyltransferase\*\*\* (LCAT) than on ACAT. In addition, it inhibited

vivo cholesterol absorption in rats by 50% at 75 mg/kg.

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3 L3 AND TRANSFORM?

=> d 15 1-3 ibib ab

L5 ANSWER 1 OF 3 BIOSIS COPYRIGHT (c) 2004 The Thomson Corporation. on STN ACCESSION NUMBER: 2004:441893 BIOSIS

DOCUMENT NUMBER:

PREV200400446784

TITLE:

in

Expression in yeast of a novel phospholipase Al cDNA from

Arabidopsis thaliana.

AUTHOR(S): Noiriel, Alexandre; Benveniste, Pierre; Banas, Antoni;

Stymne, Sten; Bouvier-Nave, Pierrette [Reprint Author]

CORPORATE SOURCE: CNRSInst Biol Mol PlantesDept Isoprenoides, Inst Bot, 28

Rue Goethe, F-67083, Strasbourg, France Pierrette.Nave@bota-ulp.u-strasbg.fr

SOURCE: Furopean Journal of Riochemistry (9

European Journal of Biochemistry, (September 2004) Vol.

271, No. 18, pp. 3752-3764. print. ISSN: 0014-2956 (ISSN print).

DOCUMENT TYPE: Article

LANGUAGE:

English

ENTRY DATE:

Entered STN: 17 Nov 2004

Last Updated on STN: 17 Nov 2004

AB During a search for cDNAs encoding \*\*\*plant\*\*\* sterol acyltransferases, we isolated four full-length cDNAs from Arabidopsis thaliana that encode proteins with substantial identity with animal \*\*\*lecithin\*\*\* : \*\*\*cholesterol\*\*\* \*\*\*acyltransferases\*\*\* (LCATs). The expression of one of these cDNAs, AtLCAT3 (At3g03310), in various yeast strains resulted in the doubling of the triacylglycerol

content. Furthermore, a complete lipid analysis of the \*\*\*transformed\*\*\* wild-type yeast showed that its phospholipid content was lower than that of the control (void plasmid- \*\*\*transformed\*\*\* ) yeast whereas lysophospholipids and free fatty acids increased. microsomes from the AtLCAT3- \*\*\*transformed\*\*\* yeast were incubated with di-(1-14C)oleyl phosphatidylcholine, both the lysophospholipid and free fatty acid fractions were highly and similarly labelled, whereas the same incubation with microsomes from the control yeast produced a negligible labelling of these fractions. Moreover when microsomes from AtLCAT3- \*\*\*transformed\*\*\* yeast were incubated with either sn-1- or sn-2-(1-14C)acyl phosphatidylcholine, the distribution of the labelling between the free fatty acid and the lysophosphatidylcholine fractions strongly suggested a phospholipase A1 activity for AtLCAT3. The sn-1 specificity of this phospholipase was confirmed by gas chromatography analysis of the hydrolysis of 1-myristoyl, 2-oleyl phosphatidylcholine. Phosphatidylethanolamine and phosphatidic acid were shown to be also hydrolysed by AtLCAT3, although less efficiently than phosphatidylcholine. Lysophospatidylcholine was a weak substrate whereas tripalmitoylglycerol and cholesteryl oleate were not hydrolysed at all. This novel A. thaliana phospholipase Al shows optimal activity at pH 6-6.5 and 60-65 degreeC and appears to be unaffected by Ca2+. Its sequence is unrelated to all other known phospholipases. Further studies are in progress to elucidate its physiological role.

L5 ANSWER 2 OF 3 CAPLUS COPYRIGHT 2004 ACS on STN

ACCESSION NUMBER:

2001:168132 CAPLUS

DOCUMENT NUMBER:

134:218021

TITLE:

Nucleic acids encoding \*\*\*plant\*\*\* sterol acyltransferases and their use to modify sterol

composition

INVENTOR(S):

Lassner, Michael; Van Eenennaam, Alison

PATENT ASSIGNEE(S): SOURCE:

Monsanto Company, USA

PCT Int. Appl., 127 pp. CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

```
KIND
    PATENT NO.
                              DATE
                                         APPLICATION NO.
                                                               DATE
                       ----
                              -----
                                         _______
    WO 2001016308
                       A2
                              20010308
                                         WO 2000-US23863
     WO 2001016308
                       A3
                              20020117
        W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,
            CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HR,
            HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT,
            LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU,
            SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN,
            YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
        RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY,
            DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ,
            CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
    CA 2381901
                        AA 20010308 CA 2000-2381901
                                                               20000830
    BR 2000014154
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    EP 1210417
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            IE, SI, LT, LV, FI, RO, MK, CY, AL
    JP 2003508052
                        T2
                              20030304
                                         JP 2001-520855
                                                               20000830
     ZA 2002001410
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                              20030606
                                        ZA 2002-1410
                                                               20020219
PRIORITY APPLN. INFO.:
                                         US 1999-152493P
                                                            P 19990830
                                         WO 2000-US23863
                                                            W 20000830
AΒ
    The present invention is directed to
                                         ***lecithin*** :
      and
    acyl CoA: cholesterol acyltransferases-like polypeptides (ACAT).
    invention provides polynucleotides encoding such
    cholesterol:acyltransferase-like polypeptides, polypeptides encoded by
    such polynucleotides, and the use of such polynucleotides to alter sterol
    compn. and oil prodn. in ***plants*** and host cells. Four LCAT cDNAs
    are provided from Arabidopsis thaliana, as well as 2 genomic DNAs encoding
    LCAT from A. thaliana, 7 ESTs from soybean and 11 ESTs from corn.
    ACAT-encoding ESTs are also identified from A. thaliana, soybean, maize,
    and Mortierella alpina. Also provided are oils produced by the
                    and host cells contq. the polynucleotides and food
    products, nutritional supplements, and pharmaceutical compn. contq.
                    or oils of the present invention.
L5
    ANSWER 3 OF 3 CAPLUS COPYRIGHT 2004 ACS on STN
ACCESSION NUMBER:
                       2000:384442 CAPLUS
DOCUMENT NUMBER:
                       133:27387
TITLE:
                       Polynucleotides (cDNA) and polypeptides of
                         ***acyltransferase***
                                              sequence homologs, sequences
                       and biological uses thereof
INVENTOR (S):
                       Cahoon, Rebecca E.; Kinney, Anthony J.; Sakai, Hajime;
                       Shen, Jennie Bih-jien; Butler, Karlene H.; Saylor,
PATENT ASSIGNEE(S):
                       E. I. Du Pont de Nemours & Co., USA
SOURCE:
                       PCT Int. Appl., 49 pp.
                       CODEN: PIXXD2
DOCUMENT TYPE:
                       Patent
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English

LANGUAGE:

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

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PATENT NO.
                        KIND
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    WO 2000032791
                         A2
                               20000608
                                          WO 1999-US28586
    WO 2000032791
                         Α3
                               20000914
        W: AE, AL, AU, BA, BB, BG, BR, CA, CN, CR, CU, CZ, DM, EE, GD, GE,
            HR, HU, ID, IL, IN, IS, JP, KP, KR, LC, LK, LR, LT, LV, MG, MK,
            MN, MX, NO, NZ, PL, RO, SG, SI, SK, SL, TR, TT, UA, US, UZ, VN,
            YU, ZA, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
        RW: GH, GM, KE, LS, MW, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE,
            DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF,
            CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
PRIORITY APPLN. INFO.:
                                          US 1998-110782P
    The invention provides cDNA mols. encoding corn and soybean
      ***lecithin*** : ***cholesterol***
                                              ***acyltransferases***
    based on sequence homol. to known LCATs. The invention also provides a
    chimeric gene comprising the ***plant***
                                                LCAT cDNA operably linked to
    suitable regulatory sequences (such as promoter and terminator sequences)
    and a host cell (such as yeast, bacteria,
                                              ***plant***
                                                             or virus)
      ***transformed***
                         with said chimeric gene for the recombinant prodn. of
    the LCAT. The invention further provides for the use of: (1)
      ***plant***
                    LCAT-specific primers for amplification of a nucleic acid
    encoding LCAT; (2)
                        ***plant*** LCAT-specific probes in screening a
    cDNA or genomic library for nucleic acid mols. encoding LCAT and (3)
    polynucleotides comprising at least 30 nucleotides of the LCAT cDNA mol.
    or complement of such sequence, used for identifying an polynucleotide
    that affects the level of LCAT expression. Finally, the invention
    provides: (1) a method for evaluating the ability of a mol. to inhibit the
    activity of LCAT and (2) a method for selecting ***transformed***
      ***plant*** cells overexpressing LCAT, which involves measuring the
    phytosterol concn. in the cell. CDNA and amino acid sequences of full
    length and partial cDNA clones encoding the corn LCAT sequence homologs
    are provided. Likewise, cDNA and amino acid sequences of a full length
    and a partial cDNA clone encoding soybean LCAT sequence homologs are
    provided. Using the BLASTX algorithm, the amino acid sequences of various
    putative corn LCATs were found to be 29.4% to 37.2% similar to the amino
    acid sequence of Arabidopsis thaliana GenBank accession no. AC004557
    GI3935185, while the sequence of the putative soybean LCAT was found to be
    57.% similar to the sequence of A. thaliana. The invention also discussed
    that overexpression or cosuppression of LCAT may be useful to genetically
    alter the content of phytosterol or lecithin in grains.
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| COST IN U.S. DOLLARS                       | SINCE FILE | TOTAL   |
|  | ENTRY      | SESSION |
| FULL ESTIMATED COST                        | 90.35      | 90.56   |
|  |            |         |
| DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS) | SINCE FILE | TOTAL   |
|  | ENTRY      | SESSION |
| CA SUBSCRIBER PRICE                        | -2.10      | -2.10   |

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CA SUBSCRIBER PRICE 0.00 -2.10

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ANSWER 26 OF 64 BIOSIS COPYRIGHT (c) 2004 The Thomson Corporation.

ACCESSION NUMBER: 1995:24979 BIOSIS DOCUMENT NUMBER: PREV199598039279

TITLE: Oyster mushroom (Pleurotus ostreatus) decreases serum and

liver cholesterol and increases cholesterol

7-alpha-hydroxylase activity and fecal excretion of neutral

sterols and bile acids in hypercholesterolemic rats.

AUTHOR(S): Bobek, P.; Ondreicka, R.; Klvanova, J.; Ozdin, L. Res. Inst. Nutrition, Limbova 14, Bratislava 833 37, CORPORATE SOURCE:

Slovakia

SOURCE: Nutrition Research, (1994) Vol. 14, No. 11, pp. 1683-1688.

CODEN: NTRSDC. ISSN: 0271-5317.

DOCUMENT TYPE: Article LANGUAGE: English

ENTRY DATE: Entered STN: 11 Jan 1995

Last Updated on STN: 12 Jan 1995

AB Wistar rats fed a semisynthetic diet containing 0.3% cholesterol and supplemented with 5% dried whole oyster mushroom (Pleurotus ostreatus) had reduced serum and liver cholesterol levels by 32 and 55%, respectively, at the end of 8th week of the experiment. The reduction of cholesterol was due to the decreased cholesterol content in very-low-density lipoproteins (VLDL) and in low-density lipoproteins (LDL). Cholesterol concentration in high-density lipoproteins (HDL) increased significantly by 34%. Animals fed the oyster mushroom diet had elevated level of fecal excretion of neutral sterols by 32% and the excretion of bile acids by 55%. Activity of cholesterol 7-alpha-hydroxylase (a rate-limiting enzyme of cholesterol catabolism) was enhanced by 33% and the activity of : \*\*\*cholesterol\*\*\* \*\*\*lecithin\*\*\* \*\*\*acyltransferase\*\*\*

increased by 13%.

ANSWER 27 OF 64 EMBASE COPYRIGHT 2004 ELSEVIER INC. ALL RIGHTS RESERVED.

on STN

ACCESSION NUMBER: 94358700 EMBASE

DOCUMENT NUMBER:

1994358700

TITLE: Hypolipedaemic activity of picroliv in albino rats. AUTHOR: Khanna A.K.; Chander R.; Kapoor N.K.; Dhawan B.N.

CORPORATE SOURCE:

Division of Biochemistry, Central Drug Research Institute,

PO Box 173, Lucknow 226001, India

SOURCE:

Phytotherapy Research, (1994) 8/7 (403-407).

ISSN: 0951-418X CODEN: PHYREH

COUNTRY:

United Kingdom

DOCUMENT TYPE:

Journal; Article 018

FILE SEGMENT:

Cardiovascular Diseases and Cardiovascular Surgery

Clinical Biochemistry 029

030 Pharmacology

037 Drug Literature Index

LANGUAGE:

English SUMMARY LANGUAGE: English

The hypolipidaemic action of picroliv, a standarized preparation from Picrorhiza kurrooa, has been studied in normal as well as in triton- and cholesterol-fed rats. Serum lipids were found to be lowered by picroliv (25 mg/kg b.w.) in triton WR-1339-induced hyperlipaemia. Chronic feeding of this drug (6 mg/kg b.w.) in normal rats and in animals simultaneously treated with cholesterol (25 mg/kg b.w.) for 30 days caused lowering in the lipid and protein levels constituting .beta.-lipoproteins followed by an increase in high density lipoprotein cholesterol in experimental animals. Picroliv alters lipolytic activities in plasma, liver, heart an adipose tissues and stimulated receptor mediated catabolism of low density lipoprotein. The lipid lowering action of the natural product is mediated through inhibition of cholesterol biosynthesis in liver, increased faecal bile acid excretion and enhanced plasma \*\*\*lecithin\*\*\* \*\*\*cholesterol\*\*\* \*\*\*acyltransferase\*\*\* activity.

ANSWER 28 OF 64 BIOSIS COPYRIGHT (c) 2004 The Thomson Corporation. 1.3 STN

ACCESSION NUMBER: DOCUMENT NUMBER:

1993:367422 BIOSIS PREV199396053097

TITLE:

The role of cholesterol absorption and hepatic cholesterol content in high and low responses to dietary cholesterol

and fat in pedigreed baboons (Papio species).

AUTHOR (S):

SOURCE:

Kushwaha, Rampratap S. [Reprint author]; Rice, Karen S.; Lewis, Douglas S.; McGill., Henry C., Jr.; Carey, K. D.

CORPORATE SOURCE:

Dep. Physiol. and Med., Southwest Foundation Biomedical Res., PO Box 28147, San Antonio, TX 78228-0147, USA

Metabolism Clinical and Experimental, (1993) Vol. 42, No.

6, pp. 714-722.

CODEN: METAAJ. ISSN: 0026-0495.

DOCUMENT TYPE:

Article English

LANGUAGE: ENTRY DATE:

Entered STN: 6 Aug 1993

Last Updated on STN: 8 Aug 1993

Selective breeding has produced baboon families with low and high plasma AB cholesterol responses to dietary cholesterol and fat. We used 12 highand 12 low-responding (mainly in low-density lipoprotein (LDL) cholesterol) pedigreed baboons to determine whether cholesterol absorption and hepatic cholesterol concentration are associated with these responses. We measured cholesterol absorption first on the chow diet, which was low

in cholesterol and fat, and after 3 and 13 weeks on the challenge diets, which contained 0.45 mg cholesterol/kcal and 40% of calories as either coconut oil or corn oil. Plasma, lipoprotein, and hepatic cholesterol concentrations were measured 1 week after cholesterol absorption measurements. High-responding baboons had higher percentage cholesterol absorption than low-responding baboons on both chow and challenge diets, regardless of the type of dietary fat. Both high and low responders had higher percentage cholesterol absorption with corn oil than with coconut oil. High responders also had higher hepatic cholesterol concentrations than low responders on chow and after consuming the challenge diets for 4 weeks. After consuming the challenge diets for 14 weeks, low responders fed coconut oil had hepatic cholesterol levels equal to those of high responders, while low responders fed corn oil continued to have low hepatic cholesterol levels. Thus, percentage cholesterol absorption is consistently higher in high-responding baboons regardless of diet, but hepatic cholesterol concentration varies with duration of challenge and type of fat. The results suggest that both cholesterol absorption and hepatic cholesterol concentration regulate cholesterolemic responses to diet, but by different mechanisms.

 $L_3$ ANSWER 29 OF 64 EMBASE COPYRIGHT 2004 ELSEVIER INC. ALL RIGHTS RESERVED.

on STN

ACCESSION NUMBER: 92313699 EMBASE

DOCUMENT NUMBER:

1992313699

TITLE:

The effect of borage oil consumption on human plasma lipid

levels and the phosphatidylcholine and cholesterol ester

composition of high density lipoprotein.

AUTHOR:

Barre D.E.; Holub B.J.

CORPORATE SOURCE:

Department of Nutritional Sciences, University of

Guelph, Guelph, Ont. N1G 2W1, Canada

SOURCE:

Nutrition Research, (1992) 12/10 (1181-1194).

ISSN: 0271-5317 CODEN: NTRSDC

COUNTRY:

United States

DOCUMENT TYPE:

Journal; Article

FILE SEGMENT:

017 Public Health, Social Medicine and Epidemiology

029 Clinical Biochemistry 037 Drug Literature Index

LANGUAGE:

English

SUMMARY LANGUAGE: English

The effect of consuming an enriched source of gamma-linolenic acid (GLA, AΒ 18:3 n-6) in the form of borage oil on the plasma lipid levels of human volunteers and the fatty acid composition of high density lipoprotein (HDL) - phosphatidylcholine (PC) and HDL- cholesterol ester (CE) was examined. Furthermore, an estimation was made of the relative utilization of GLA and dihomo-gamma-linolenic acid (DGLA, 20:3 n-6) by the plasma \*\*\*lecithin\*\*\* : \*\*\*cholesterol\*\*\* \*\*\*acyltransferase\*\*\* reaction. For this purpose, six healthy male subjects consumed encapsulated borage oil (21.8 wt % GLA) so as to provide an average intake of 5.23 g GLA/person/day for 42 consecutive days followed by a wash-out period of an additional 42 days. Analysis of plasma lipids (triglycerides, total cholesterol, HDL-cholesterol, low density lipoprotein (LDL) cholesterol) and the plasma HDL-cholesterol:total cholesterol ratio indicated no significant change in any of the latter arising from the supplementation. Consumption of the borage oil supplement was found to produce marked alterations in the fatty acid compositions of HDL-PC and HDL-CE. Whereas only a moderate accumulation in PC (up to 0.6 mol %) of GLA was found with supplementation, a more dramatic rise in DGLA (by 3.9

mol %) and arachidonic acid (AA, 20:4 n-6) (by 3.3 mol %) was found. These changes were reversible by day 64. In contrast to the HDL-PC, the GLA accumulation in HDL-CE exceeded that of DGLA such that the net rise was 1.8 mol % as compared to only 0.8 mol % for DGLA. Positional analysis of the HDL- PC from subjects at day 43 revealed GLA and DGLA to reside almost exclusively in the 2-position (> 98 %). Dividing the percentages of unsaturated fatty acids in the HDL-CE relative to their levels in the corresponding 2-position of PC provided the following estimated order of fatty acid selectivity for the plasma LCAT reaction: GLA > linoleic acid (LA, 18:2 n-6) > oleic acid (OA, 18:1 n-9) > AA > DGLA .gtoreq. docosahexaenoic acid (DHA, 22:6 n-3).

L3 ANSWER 30 OF 64 BIOSIS COPYRIGHT (c) 2004 The Thomson Corporation. on

ACCESSION NUMBER:

1993:98481 BIOSIS

DOCUMENT NUMBER:

PREV199395053677

TITLE:

Effect of dietary safflower phospholipid and soybean

phospholipid on plasma and liver lipids in rats fed a

cholesterol-free diet.

AUTHOR (S):

Iwata, Toshio [Reprint author]; Takehisa, Fumiyuki;
Tsutsumi, Kentarou; Furukawa, Yuji; Kimura, Shuichi

CORPORATE SOURCE:

Dep. Res. Development, Rinoru Oil Mills Co., Ltd.,

Minato-ku, Nagoya 455, Japan

SOURCE:

Journal of Clinical Biochemistry and Nutrition, (1992) Vol.

13, No. 2, pp. 107-115.

CODEN: JCBNER. ISSN: 0912-0009.

DOCUMENT TYPE:

Article English

LANGUAGE: ENTRY DATE:

Entered STN: 9 Feb 1993

Last Updated on STN: 10 Feb 1993

AB The effect of dietary safflower phospholipid (Saf-PL) and soybean phospholipid (Soy-PL) on plasma, lipid, and fecal lipids in rats fed a cholesterol-free diet was compared with that of a triglyceride mixture (control). The triglyceride mixture (SP-Oil) of safflower oil and palm oil (8:2) contained almost comparable amounts of linoleic acid to safflower phospholipid or soybean phospholipid. Concentrations of total cholesterol in plasma of rats fed the Saf-PL and Soy-PL diets were significantly decreased in comparison with that of the SP-Oil diet. Saf-PL induced a reduction in the concentration of liver cholesterol compared with SP-Oil. Soy-PL tended to reduce the liver cholesterol. The proportions of total cholesterol in all lipoprotein fractions were similar among the groups. The activity of plasma \*\*\*lecithin\*\*\* -

\*\*\*cholesterol\*\*\*

\*\*\*acyltransferase\*\*\*

was increased in rats fed the phospholipid diets; Saf-PL indicated the highest value. Saf-PL and Soy-PL caused an enhanced excretion of not only neutral steroids but also acidic steroids into feces compared with SP-Oil. These results suggest that, in addition to soybean phospholipid, safflower phospholipid decreases plasma and liver cholesterol in rats fed a cholesterol-free diet and safflower phospholipid causes a favorable alteration in plasma and liver lipids compared with soybean phospholipid.

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STN INTERNATIONAL LOGOFF AT 16:10:45 ON 10 DEC 2004